

Dirty appearing white matter in the brain is associated with altered cerebrospinal fluid pulsatility and hypertension in individuals without neurologic disease

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Short title: Dirty appearing white matter and CSF pulsatility

Type of manuscript: original research

Keywords: Dirty appearing white matter; cerebrospinal fluid; hypertension; leukoaraiosis; magnetic resonance imaging

Sources of funding: This work has been supported in part by a grant from the Annette Funicello Research Fund for Neurological Diseases.

Disclosures

Clive Beggs, Christopher Magnano, Simon Shepherd, Pavel Belov, Deepa Ramasamy and Jesper Hagemeier have nothing to disclose. Robert Zivadinov received personal compensation from Teva Pharmaceuticals, Biogen Idec, EMD Serono and Genzyme for speaking and consultant fees. Dr. Zivadinov also received financial support for research activities from Biogen Idec, Teva Pharmaceuticals, Genzyme and Novartis.

Abstract count: 247, Word count (text and references only): 4945, Number of Tables: 4, Number of Figures: 2, Number of references: 48.

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Abstract

Background and purpose

Aging of the healthy brain is characterized by focal or non-focal white matter (WM) signal abnormality (SA) changes, which are typically detected as leukoaraiosis (LA). Hypertension is a risk factor for WM lesion formation. This study investigated whether LA might be associated with increased CSF pulsatility linked to arterial hypertension.

Methods

101 individuals without neurologic diseases (53 females and 48 males) aged between 18-75 years underwent 3T brain MRI with cine phase contrast imaging for CSF flow estimation, after providing their informed consent. LA was defined as the presence of focal T2 WM SA changes and/or non-focal uniform areas of signal increase termed dirty-appearing-white-matter (DAWM). Relevant information relating to cardiovascular risk factors was also collected.

Results

When controlled for age and hypertension, significant partial correlations were observed between: DAWM volume and: net negative flow ($r=-0.294$, $p=0.014$); net positive flow (NPF) ($r=0.406$, $p=0.001$); and peak positive velocity ($r=0.342$, $p=0.004$). Multiple linear regression analysis revealed DAWM volume to be significantly correlated with CSF NPF ($p=0.019$) and hypertension ($p=0.007$), whereas T2 WM SA volume was only significantly correlated with age ($p=0.002$). Combined DAWM and T2 WM SA volumes were significantly related with age ($p=0.001$) and CSF peak negative velocity ($p=0.041$).

Conclusions

Rarefaction of WM leading to LA is a multifactorial process, in which formation of DAWM induced by hypertension and increased aqueductal CSF pulsatility, may play a contributory role. These two factors appear to act independently of each other in a process that is independent of age.

Introduction

Aging of the brain in healthy individuals is characterized by atrophy and focal or non-focal white matter (WM) signal abnormality (SA) changes, which are typically detected as leukoaraiosis (LA).¹ However, much remains unknown about the pathophysiology underlying these changes. WM SA changes of presumed vascular origin are a common finding in brain MRI of older individuals and contribute to cognitive and functional decline.² Although WM degeneration is characterized pathologically by demyelination, axonal loss, and rarefaction (processes often attributed to ischemia),³ it is not known precisely how WM SA changes form.⁴ A recent study, investigating the intensity patterns and morphological features of periventricular WM SA changes, found hyperintensity intensity levels, distribution, and association with risk factors and disease, to be indicative of true tissue abnormalities in old age, and something that should not be dismissed as artifacts.⁴ In particular, the role of dirty appearing white matter (DAWM) (sometimes termed 'diffusely abnormal white matter') formation in the periventricular region is poorly understood.⁵ DAWM (Figure 1) is defined as a non-focal, uniform region of intermediate signal intensity between that of focal T2 WM SA changes and that of normal-appearing WM. Image-pathology correlation studies reveal that non-focal DAWM changes, compared with focal T2 WM SA changes, are associated with mild blood-brain barrier breakdown and myelin loss.⁶ Therefore, it is thought that non-focal DAWM may represent an early stage of WM pathology,⁶ which is followed at a later stage by focal T2 WM SA formation.^{7,8}

Recently we undertook a study involving multiple sclerosis (MS) patients,⁹ in which we found WM lesion formation to be associated with increased cerebrospinal fluid (CSF) pulsatility in the aqueduct of Sylvius (AoS). Follow-up work involving healthy subjects linked increased aqueductal pulsatility with biomechanical changes in the intracranial space associated with altered cerebral venous hemodynamics.¹⁰ In addition, we found heart disease, overweight and smoking to be associated with increased prevalence of extracranial venous abnormalities,¹¹ suggesting that cardiovascular risk factors might also influence pulsatility within the intracranial space.

It has been postulated that the presence of cardiovascular comorbidities, in particular arterial hypertension (a known risk factor for WM SA formation)¹² might be linked to increased CSF pulsatility.¹³ Hypertension is associated with reduced vascular compliance,¹⁴ particularly in smaller arterial vessels.¹⁵ This tends to promote increased pulsatility in the cerebral vascular bed,^{16,17} something that is thought to cause endothelial damage and luminal change.¹⁴ Increased cerebral blood flow pulsatility has been found to be associated with microstructural changes in the WM.^{16,18,19} Therefore, given that increased CSF pulsatility in the AoS is thought to be associated with greater pulsation in the cerebral vascular bed,²⁰ it can be

postulated that increased aqueductal CSF pulsatility might also be associated with DAWM formation.

We therefore hypothesized that DAWM formation and WM SA changes in the brain might be associated with increased aqueductal CSF pulsatility linked to arterial hypertension. In order to test this hypothesis, we undertook a study involving 101 individuals without neurologic diseases with the aim of establishing whether or not WM SA changes, and in particular the presence of DAWM, are associated with increased aqueductal pulsatility linked to cardiovascular comorbidities.

Methods

Patient population

This study utilized data from an ongoing prospective study of genetic and environmental risk factors in individuals with and without neurologic diseases that enrolled over 1,100 subjects.^{21,22} One hundred and one consecutive individuals without neurologic diseases (53 females and 48 males) aged between 18 to 75 years who underwent magnetic resonance imaging (MRI) scan with cine phase contrast imaging for CSF flow estimation were included. The individuals needed to qualify on a health screening questionnaire containing information about medical history (illnesses, surgeries, medications, etc.) and meet the health screen requirements for MRI on physical examination, as previously described.^{22,23,24} Exclusion criteria were: pre-existing medical conditions known to be associated with brain pathology (e.g. cerebrovascular disease, positive history of alcohol abuse, etc.), history of cerebral congenital vascular malformations, or pregnancy. Relevant information relating to cardiovascular risk factors [body mass index (BMI), hypertension, hypercholesterolemia, heart disease, diabetes and smoking] was collected.

All participants underwent clinical and MRI examinations. The study was approved by the University of Buffalo Institutional Review Board and written informed consent was obtained from all subjects.

MRI acquisition and analysis

The subject's brains were examined on a 3 Tesla GE Signa Excite HD 12.0 Twin Speed scanner (General Electric, Milwaukee, WI). All sequences were run on an 8-channel head and neck (HDNV) coil. All analyses were performed in a blinded manner.

Brain MRI sequences included 3D T1-weighted imaging (WI) using a fast spoiled gradient echo (FSPGR) with magnetization-prepared inversion recovery (IR) pulse for segmentation;

cine phase contrast imaging for CSF flow estimation; multi-planar dual fast spin-echo proton density (PD) and T2-WI; and fluid attenuated inversion recovery (FLAIR) for lesion analysis. A sagittal T2-weighted fast SE sequence was also acquired as a localizer for the cine phase contrast (PC) prescription, as previously described, with the cine PC sequence prescribed as an oblique axial slice through the AoS.⁹

Pulse sequence characteristics for 3T MRI were as follows: all scans were acquired with a 256 x 256 matrix and a 25.6 cm FOV for an in-plane resolution of 1 x 1 mm² with a phase FOV of 75% and one average. Sequence-specific parameters were as follows: for 3D HIRIS, T1-WI were Echo Time/Inversion Time/Repetition Time TE/TI/TR=2.8/900/5.9 ms, Flip Angle (FA)=10°, and 1mm³ isotropic voxels; for the PD/T2: 3-mm-thick slices with no gap, TE1/TE2/TR = 12/95/3000 ms, ETL = 14, and for the FLAIR scans, 3-mm-thick slices with no gap, TE/TI/TR=120/2100/8500 ms.

CSF flow quantification was performed using a single slice cine phase-contrast velocity-encoded pulse-gated gradient echo sequence (cine PC) with an TE/TR of 7.9/40 ms, a slice thickness of 4 mm, a velocity encoding of 20 cm/s and 32 phases acquired corresponding to the cardiac cycle. All subjects underwent the MRI exam during the same time of day (in the afternoon hours) to control for circadian variation. The cine PC sequence was acquired with a resolution of 0.39 x 0.39 x 4 mm³, with the AoS in the center of the FOV, such that the wrap around artifact was present in the edges of the FOV, but did not overlap with the desired region of interest (ROI).

Cine phase contrast image analysis

The net positive and net negative flows (NPF and NNF), together with the net flow (NF = NNF + NPF) and the peak positive and negative velocities (PPV and PNV) were calculated using a validated methodology as previously described.⁹ The CSF flow data was processed using GE ReportCard software (version 3.6; General Electric, GE, Milwaukee, WI) and positive and negative velocities over all 32 phases were recorded. In order to overcome limitations in the ReportCard software an in-house semi-automated minimum area of contour change (MACC) program was used to correct the ROIs for each phase, as previously described.⁹ NPF and NNF were calculated using only the phases, which have positive and negative velocities, respectively.⁹ CSF flow measures are presented in microliters per beat (μL/beat, 1μL = 1mm³), while CSF velocity measures are presented in cm/s. CSF flow direction was calculated based on slice prescription such that flow through the AoS out of the slice (during diastole, towards the third ventricle) was given as positive, whereas flow into the slice (during systole, towards the fourth ventricle) was negative, as described previously.⁹

Detection of white matter signal abnormality changes:

LA was defined as the presence of focal T2 WM SA changes and/or non-focal uniform areas of signal increase termed DAWM. Each subject's relevant T2/PD images were co-registered to the FLAIR using a rigid body (6-degrees-of-freedom) registration.²⁵ The FLAIR image was used to outline T2 WM SAs, while the co-registered T2/PD images were used to confirm the presence of T2 WM SAs and to check for infratentorial T2 WM SAs, as FLAIR imaging is known to be insensitive in this area. The T2 WM SA number and volume (T2 WM-SAV) were outlined using a semi-automated edge detection contouring/thresholding technique as described previously.²⁶ All T2 WM SAs were divided into individual ROIs $\geq 3\text{mm}$ in size (equivalent to $\geq 14.1\text{ mm}^3$). The periventricular and deep WM regional localization of T2 WM SAs was also determined.

In addition, areas of DAWM were outlined (Figure 1). The DAWM was defined as a uniform, non-focal area of signal increase on the FLAIR/T2/PD-weighted sequence at 3T, with a subtly increased signal intensity compared with the contralateral signal intensity of normal appearing WM but less than that of T2 WM SAs, as previously proposed.²⁷

Statistical analysis

Analysis was undertaken using in-house algorithms written in R (open source statistical software) and Matlab (Mathworks, Natick, Mass). A one-way ANOVA was used to analyze changes in CSF pulsatility with age, with values of $p < 0.05$ using a two-tailed test considered statistically significant.

Correlation (full and partial) matrices (Spearman's r) were computed to quantify the relationships between the variables. Multiple linear regression analysis using the dependent variables T2 WM-SAV, DAWM volume, DAWM volume+T2 WM-SAV, T2 periventricular SA number (T2 PVL-SAN) and T2 deep WM SA number (T2 DWM-SAN) was also performed. In this analysis the independent variables were age, BMI, hypertension, hypercholesterolemia, heart disease, diabetes, smoking, NF, NNF, NPF, PPV and PNV. A stepwise approach was taken, with variables excluded according to their respective t values. Only significant variables with $p < 0.05$ were included in the final regression models.

Values of $p < 0.05$ using a two-tailed test were considered statistically significant after the Benjamini-Hochberg²⁸ correction for multiple comparisons was applied.

Results

Demographic and clinical characteristics

Table 1 shows the demographic, clinical and MRI characteristics. Subject average age was 44.7 years (range: 18 – 75 years), with females comprising 54.5% of the cohort. Recorded cardiovascular risk factors were as follows: 8.2% had a history of cardiovascular disease; 22.8% were smokers; 27.8% had hypertension; 17.0% had hypercholesterolemia; with 6.2% having diabetes. The mean BMI was 26.8 kg/m². With the exception of smoking, no significant differences were observed between the male and female subjects. The MRI characteristics are also presented in Table 1.

One-way ANOVA of the CSF data grouped according to age (Figure 2), revealed statistical differences for NF ($p=0.003$), NPF ($p=0.021$) and NPV ($p=0.030$), suggesting that the values of these variables changed with age.

Correlation analysis

Correlation analysis of the demographic and cardiovascular risk factor variables against the MRI variables (Table 2) revealed age to be significantly correlated with NF, PNV, T2 WM-SAV, DAWM volume and DAWM volume+T2 WM-SAV, while hypertension was significantly correlated with T2 WM-SAV, DAWM volume and DAWM volume+T2 WM-SAV, with BMI significantly correlated with T2 WM-SAV.

Correlation analysis of the relationships between the MRI variables (Table 3a) revealed significant correlations between the CSF variables NNF, NPF, PPV and PNV, and the MRI variables T2 WM-SAV, DAWM volume and DAWM volume+T2 WM-SAV. Significant observations were also observed between: T2 DWM-SAN and the variables NNF, NPF and PPV; and T2 PVL-SAN and NPF.

Partial correlation analysis

When controlled for age and hypertension, the partial correlation analysis (Table 3b) revealed significant correlations between: DAWM volume and NNF, NPF and PPV; DAWM volume+T2 WM-SAV and NPF and PPV; and T2 DWM-SAN and NNF, NPF.

Linear regression analysis

Multiple linear regression analysis was undertaken with T2-SAV, DAWM volume, DAWM volume+T2 WM-SAV, T2 PVL-SAN and T2 DWM-SAN as dependent variables. The results of this analysis (Table 4) revealed that significant models were only achieved for T2 WM-SAV, DAWM volume, and the composite variable DAWM volume+T2 WM-SAV. Interestingly, there were marked differences between the models for T2 WM-SAV and DAWM volume. For

the dependent variable T2 WM-SAV, the only significant correlation was age, whereas for DAWM volume, hypertension and NPF were significant. The results for the DAWM volume+T2 WM-SAV model reflected the composite nature of that variable, with age and PNV both being significant.

Discussion

Our principal finding is that in individuals without neurologic diseases, DAWM formation appears to be associated with both hypertension and increased aqueductal CSF pulsatility, rather than aging *per se*. When controlled for age, the partial correlations between DAWM volume and the CSF variables, NNF, NPF, PPV and PNV were all significant, with the partial correlation between DAWM volume and hypertension approaching significance. This suggests that rarefaction of the WM leading to LA is a multifactorial process, in which formation of DAWM is influenced by factors other than age. As such, our findings mirror those of Jolly *et al.*¹⁹ who found microstructural WM changes to be independent of age and associated with increased cerebral blood flow pulsatility and aqueductal CSF pulse volume.

Cerebral WM SA changes frequently occur in healthy ageing and are typically detected as LA, a radiological finding that presents as increased signal intensity on T2-weighted MR images. WM SA changes are a common finding in brain MRI of older individuals and are thought to contribute to cognitive and functional decline.² While it is not known precisely how these WM SA changes form, it has been shown that periventricular WM hyperintensities in old age represent true tissue abnormalities.⁴ WM degeneration is characterized pathologically by demyelination, axonal loss, and rarefaction, which are thought to be related to ischemia.³ Histological studies have shown LA to be characterized by WM morphological changes around the periventricular veins.^{29,30} In particular, LA is associated with non-inflammatory collagenosis of the periventricular veins, resulting in thickening of the vessel walls and narrowing of the lumen.²⁹ Vascular dementia, closely related to LA, has been shown to be associated with alterations in venous pulsation,³¹ and it has been postulated that the thickening of the vein walls in LA may be a protective response against increased vascular pulsatility.²⁹ Other subtle changes appear to be associated with increased vascular pulsatility. Jolly *et al.*¹⁹ found fractional diffusion anisotropy to be negatively correlated with both arterial pulse and pulsatility in deep venous territories. Changes in diffusion anisotropy have also been reported in the normal-appearing white matter around LA lesions³² and it is thought that these represent early changes in the structural organization of white matter that are likely to precede the emergence of LA.¹⁹

DAWM is poorly understood and appears something of an enigma. Most of the work undertaken on DAWM has been done in the context of MS, where it has been shown that quantitative MR imaging measures of DAWM lie between those of normal-appearing WM and focal WM lesions.^{33,34} This has led some to suggest that DAWM may represent an early stage of WM pathology,⁶ which is followed at a later stage by focal WM lesion formation. In support of this, Chung *et al.*³⁵ reported that DAWM appears to be converted to T2 WM lesions in Alzheimer's disease (AD) patients who experience severe jugular venous reflux. Others however, suggest that DAWM is by no means a preceding stage for focal lesion formation, but rather a different pathologic phenomenon altogether, arising from secondary Wallerian degradation and resulting in extensive axonal and myelin loss.^{34,36}

While the pathology associated with DAWM remains unclear, there appears to be a consensus that DAWM reflects an increase in extracellular water within the brain parenchymal tissue.³³ The apparent diffusion coefficient (ADC) has been shown to be greater in DAWM compared with normal-appearing WM in patients with MS,³⁴ indicating the presence of microstructural changes in DAWM that permit increased transport of water. Laule *et al.*³⁷ found that the primary abnormality in DAWM was a reduction or perturbation of the myelin phospholipids that was associated with a reduction in the myelin water fraction. This was accompanied by an increase in the total water content of the DAWM, probably due to an influx of water into spaces formerly occupied by the phospholipids and myelin proteins. As such, this suggests that profound microstructural changes may occur within DAWM, possibly due to lipid abnormalities.³⁸ While DAWM has been studied in MS, it has rarely been investigated outside of this context, and so it cannot be assumed that DAWM formation in healthy individuals necessarily represents the same pathophysiology as that observed in MS patients. Notwithstanding this, given that water has a long T2 value, it is reasonable to postulate that DAWM formation might also be associated with increased ADC in healthy subjects.

While the specific impact of altered CSF pulsatility on DAWM formation has not previously been investigated, a number of studies have linked increased intracranial fluid pulsatility and microstructural WM damage.^{16,18,19} Our finding that DAWM formation is associated with hypertension appears to be consistent with those who linked microstructural WM damage with increased cerebral blood flow pulsatility.^{16,18,19} Increased pulsatility is indicative of decreased arterial compliance, something that is associated with arteriosclerosis³⁹ and hypertension.¹⁴ Hypertension is a risk factor for small vessel disease⁴⁰ and LA,⁴¹ and is associated with changes in vascular mechanics.^{14,16} It has been suggested that greater vascular pulsatility might cause WM damage by increasing perivascular shear stress and

inducing demyelination, resulting in microstructural WM changes and contributing to the proliferation of LA over time.¹⁹ Stiffening of the aorta has been linked to the transmission of excessive flow pulsatility into the brain,^{16,17} something that is associated with WM microstructural changes in healthy older individuals.¹⁹ Recently, Tarumi *et al.*¹⁷ demonstrated that arterial stiffness in aging is positively correlated with cerebral vascular pulsatility, and that this in turn is associated with a greater volume of WM hyperintensities. Excessive intracranial cardiac-related pulsatility (cerebral arterial and CSF) has also been associated with brain atrophy among elderly individuals.¹⁸ Microstructural changes associated with increased cerebral pulsatility may therefore represent early stage alterations in the structural organization of the WM, likely to precede the emergence of LA.¹⁹ In our linear regression model the predictors NPF and hypertension were only able to account for about 20% of the variance associated with DAWM formation. This suggests that although both increased aqueductal pulsatility and hypertension are associated with DAWM formation, other unknown contributory factors must be at work.

The intracranial CSF windkessel mechanism dampens the vascular pulse so that blood flow through the cerebral capillary bed is non-pulsatile.⁴² If however this mechanism becomes dysfunctional, then this might lead to increased pulsatility in the AoS, as postulated by Greitz,²⁰ who argued that pulsations in the cerebral capillaries were transmitted through the parenchyma to the lateral ventricles. If this were the case, then one might expect aqueductal pulsatility to be correlated with hypertension. But when we controlled for age, we did not find any significant correlations between hypertension and the CSF variables, suggesting that the two phenomena are not linked. As such, our finding supports Rashid *et al.*⁴³ who found in a rat study that elevated pulsatile CSF flow in the aqueduct was not matched by an increase in microvascular pulsatile flow. Furthermore, our regression analysis model indicated that both hypertension and NPF were significant predictors of DAWM formation, suggesting that the two mechanisms may be acting independently of each other to produce WM SA changes. Increased aqueductal pulsatility has been linked with constricted cerebral venous outflow in healthy individuals¹⁰ and MS patients,⁴⁴ suggesting that this phenomenon may be more associated with altered compliance in the sub-arachnoid space, rather than pulsatility in the cerebral vascular bed. Notwithstanding this, a number of researchers have observed a link between the aqueductal pulsatility and subtle changes in the brain parenchyma. Daouk *et al.*⁴⁵ found ADC, an early indicator of microstructural changes, to be strongly correlated with aqueductal stroke volume in AD, and Jolly *et al.*¹⁹ found increased aqueductal CSF pulse volume to be associated with microstructural WM changes in elderly subjects. Having said this, these correlations appear to be confined microstructural changes, with Jolly *et al.* finding no association between CSF pulse volume and LA severity.¹⁹

The results regarding T2 WM SA formation appear more ambiguous than those for DAWM formation. The regression analysis model for T2 WM-SAV identified age as the only significant predictor, while those for T2 PVL-SAN and T2 DWM-SAN revealed no significant predictors. However, care should be taken before completely ruling out other contributory factors to T2 WM SA formation. This is because the NPF and NF characteristics of the >70 age group were markedly different to those <70 years of age (Figure 2). When we repeated the multiple regression analysis using only those subjects <70 years of age, we found T2 WM-SAV to be significantly correlated with hypertension rather than age (statistical power = 80.9%). Also, in the younger cohort, DAWM volume+T2 WM-SAV was significantly correlated with hypertension and NPF (statistical power = 99.2%), and DAWM volume was significantly correlated with hypertension and NPF (statistical power = 99.7%). This suggests that hypertension may be influential in both T2 WM SA and DAWM formation in young and middle-aged adults. As a risk factor for both small vessel disease⁴⁰ and LA,⁴¹ it is perhaps not surprising that we found hypertension associated with T2 WM-SA and DAWM formation. LA is characterized by thickening and hardening of vessel walls.⁴⁶ This in theory should cause blood flow in the cerebral vascular bed to become more pulsatile.^{16,17} Bateman⁴² found blood flow through the WM to be highly pulsatile in individuals with LA and concluded that this would increase endothelial shear stress, and in turn cause WM damage.³¹ Luminal narrowing associated with LA may also influence cerebral vascular density. LA is associated with decreased cerebral blood flow and capillary loss in both the cortex and the normally appearing WM.⁴⁶ This suggests that the condition affects the brain globally and that it is associated with chronic ischemia.⁴⁶ While the physiological mechanisms associated with this phenomenon are not understood, recent work by Hall *et al.*⁴⁷ has shown that ischemia associated with reduced arteriolar perfusion can result in constriction of capillaries by adjacent pericytes, rapidly leading to pericyte death and permanent occlusion of the capillaries.⁴⁸

While we found an association between aqueductal CSF pulsatility and microstructural WM changes, we did not measure the pulsatility of the cerebral blood flow. We therefore cannot assess the contribution that increased pulsatility in the cerebral vascular bed might make towards DAWM formation, or indeed, the relationship between vascular pulsatility and CSF pulsatility in the AoS.

Although we report an association between increased aqueductal pulsatility and DAWM formation in adults without neurologic diseases, previous studies have linked increased aqueductal pulsatility with: early stage WM changes¹⁹ in elderly individuals; microstructural WM changes in AD patients;⁴⁵ and increased T2 and T1 WM-SAV in MS patients.⁹ As such there appears to be an association between WM damage and aqueductal pulsatility, which is

independent of age. Therefore, we conclude that rarefaction of WM leading to LA is a multifactorial process, in which formation of DAWM induced by hypertension and increased aqueductal CSF pulsatility, may play a contributory role.

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Tables

Table 1. Descriptive statistics of the demographic, cardiovascular risk factors and MRI data.

Variables	Subjects n = 101	Males n = 46	Females n = 55	Significance p value
Age (years); mean (SD)	44.7 (17.8)	41.8 (16.1)	47.0 (19.0)	0.145
BMI (kg/m ²); mean (SD)	26.8 (5.7)	27.1 (5.0)	26.6 (6.2)	0.257
Current smokers; n (%)	23 (22.8)	15 (32.6)	8 (14.5)	0.031*
Diabetes; n (%)	6 (6.2)	2 (4.5)	4 (7.5)	0.541*
Hypertension; n (%)	20 (27.8)	5 (17.9)	15 (34.1)	0.134*
Hypercholesterolemia; n (%)	8 (17.0)	2 (12.5)	6 (19.4)	0.553*
Cardiovascular disease; n (%)	8 (8.2)	4 (8.9)	4 (7.5)	0.809*
NF (μL/beat); mean (SD)	-3.33 (7.83)	-4.35 (7.84)	-2.47 (7.79)	0.392
NNF (μL / beat); mean (SD)	-30.16 (18.73)	-31.17 (22.79)	-29.32 (14.68)	0.631
NPF (μL / beat); mean (SD)	26.83 (18.82)	26.82 (20.32)	26.85 (17.66)	0.675
PPV (cm/s); mean (SD)	6.64 (3.01)	6.87 (3.54)	6.44 (2.50)	0.865
PNV (cm/s); mean (SD)	-8.15 (3.26)	-8.32 (3.67)	-8.00 (2.91)	0.959
T2 WM-SAV; mean (SD)	350.5 (1492.7)	177.4 (646.1)	495.3 (1931.6)	0.319
DAWM volume; mean (SD)	903.1 (1113.2)	1037.0 (1355.4)	789.0 (852.0)	0.463
DAWM+T2 WM-SAV; mean (SD)	1257.1 (2049.9)	1214.4 (1785.2)	1293.5 (2267.2)	0.729
T2 PVL-SAN; mean (SD)	0.35 (1.25)	0.28 (0.78)	0.40 (1.55)	0.586
T2 DWM-SAN; mean (SD)	3.93 (8.58)	2.93 (7.92)	4.76 (9.07)	0.146

BMI, body mass index; NF, net flow; NNF, net negative flow; NPF, net positive flow; PPV, peak positive velocity; PNV, peak negative velocity; WM-SAV, white matter signal abnormality volume; DAWM, dirty appearing white matter; PVL-SAN, periventricular signal abnormality number; DWM-SAN, deep white matter signal abnormality number.

Volumes are expressed in cubic millimetres.

p value determined by 2-tailed Mann Whitney U-test unless otherwise stated.

* p value determined using chi square test.

Table 2. Spearman correlation values (r values) between demographic and cardiovascular risk factor and MRI variables (n = 101)

	NF	NNF	NPF	PPV	PNV	T2 WM-SAV	DAWM volume	DAWM+T2 WM-SAV	T2 PVL-SAN	T2 DWM-SAN
Age	0.286***^	-0.120	0.250*	0.158	-0.275***^	0.507***^	0.353***^	0.419***^	0.108	0.145
BMI	0.203	-0.120	0.217	0.061	-0.141	0.338***^	0.072	0.134	0.175	0.206
Smoker	-0.123	0.087	-0.113	-0.020	0.036	-0.024	0.071	0.075	0.005	-0.112
Diabetes	0.057	-0.147	0.182	0.122	-0.080	0.069	0.043	0.044	0.013	-0.029
Hypertension	0.185	-0.127	0.182	0.260*	-0.267*	0.485***^	0.325***^	0.353***^	0.115	0.124
Hypercholesterolemia	0.234	0.104	0.004	-0.013	-0.083	0.252	0.202	0.236	-0.189	-0.249
Cardiovascular Disease	-0.030	-0.101	0.126	0.104	-0.070	-0.025	-0.102	-0.088	-0.116	-0.055

^ Significant after Benjamini-Hochberg adjustment

*** $p \leq 0.005$ level (2-tailed).

** $p \leq 0.01$ level (2-tailed).

* $p \leq 0.05$ level (2-tailed).

Table 3. (a) Spearman correlation values (r values) between MRI white matter signal abnormality and cerebrospinal fluid flow variables; and (b) Spearman partial correlations (r values), controlling for age and hypertension, between MRI white matter signal abnormality and cerebrospinal fluid flow data. (n = 101)

		NF	NNF	NPF	PPV	PNV
(a)	T2 WM-SAV	0.201*	-0.228* [^]	0.285*** [^]	0.239* [^]	-0.264*** [^]
	DAWM volume	0.183	-0.296*** [^]	0.377*** [^]	0.343*** [^]	-0.311*** [^]
	DAWM+T2 WM-SAV	0.188	-0.290*** [^]	0.373*** [^]	0.338*** [^]	-0.318*** [^]
	T2 PVL-SAN	0.187	-0.195	0.234* [^]	0.164	-0.160
	T2 DWM-SAN	0.131	-0.290*** [^]	0.308*** [^]	0.242* [^]	-0.178
(b)	T2 WM-SAV	0.065	-0.199	0.198	0.089	-0.004
	DAWM volume	0.260*	-0.294* [^]	0.406*** [^]	0.342*** [^]	-0.268*
	DAWM+T2 WM-SAV	0.226	-0.275*	0.377*** [^]	0.309* [^]	-0.247*
	T2 PVL-SAN	0.233	-0.157	0.229	0.160	-0.131
	T2 DWM-SAN	0.107	-0.313*** [^]	0.309*** [^]	0.199	-0.063

[^] Significant after Benjamini-Hochberg adjustment

*** p ≤ 0.005 level (2-tailed).

** p ≤ 0.01 level (2-tailed).

* p ≤ 0.05 level (2-tailed).

Table 4. Results of multiple linear regression analyses for all subjects (n = 101). Dependent variables were T2 WM-SAV, DAWM volume, DAWM volume+T2 WM-SAV, T2 PVL-SAN and T2 DWM-SAN. Independent variables were age, BMI, hypertension, hypercholesterolemia, heart disease, diabetes, smoking, NF, NNF, NPF, PPV and PNv.

Dependent Variables	Independent Variables	Unstandardized Coefficients B	Unstandardized Coefficients Std. Error	Standardized Coefficients Beta	Significance p value	Adjusted r squared	Statistical power
T2 WM-SAV	Constant	-773.503	385.881		0.048	0.081	84.8%*
	Age	25.171	8.032	0.300	0.002		
DAWM volume	Constant	120.875	220.958		0.586	0.185	99.2%*
	Hypertension	794.068	284.530	0.310	0.007		
	NPF	15.923	6.610	0.268	0.019		
DAWM+T2 WM-SAV	Constant	-1395.541	620.481		0.027	0.160	97.9%*
	Age	36.778	10.956	0.321	0.001		
	PNV	-124.903	60.399	-0.198	0.041		
T2 PVL-SAN	ns	ns	ns	ns	ns	ns	na
T2 DWM-SAN	ns	ns	ns	ns	ns	ns	na

ns, not significant; na, not applicable

* Statistical power determined using post-hoc analysis

Figure Captions

Figure 1. Representation of dirty appearing white matter (DAWM – white arrow) and hyperintense T2 lesions (black arrow) in a 65 yrs old male with hypertension but not altered cerebrospinal fluid (CSF) pulsatility (upper row left) and in 69 yrs old male with altered CSF pulsatility, but no hypertension (upper row right) on fluid attenuated inversion recovery (FLAIR) sequence. The bottom row represent subjects without presence of DAWM or T2 hyperintense lesions in 64 yrs old female (left) and 64 yrs old male (right) who had no hypertension or altered CSF pulsatility.

Figures

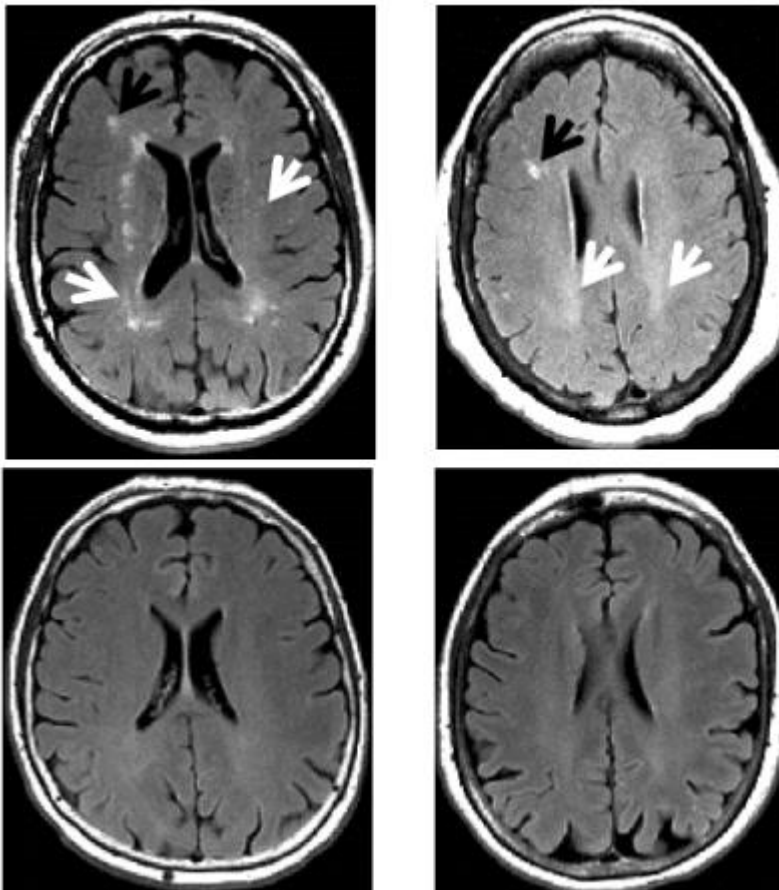


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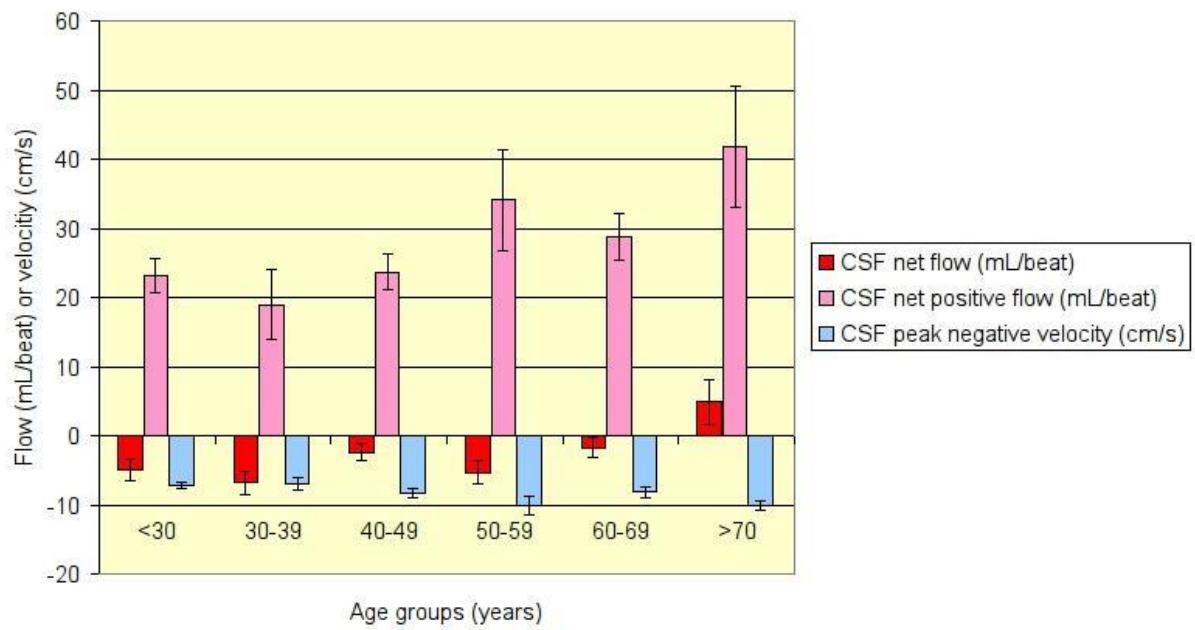


Figure 2. The effect of age on CSF NF, NPF and PNV. Error bars represent one standard error of the mean.